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- (54) Title: A NOVEL METHOD OF DIAGNOSING, MONITORING, STAGING, IMAGING AND TREATING BREAST CANCER

(57) Abstract

The present invention provides a new method for detecting, diagnosing, monitoring, staging, prognosticating, imaging and treating breast cancer.

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A NOVEL METHOD OF DIAGNOSING, MONITORING, STAGING, IMAGING AND TREATING BREAST CANCER

FIELD OF THE INVENTION

This invention relates, in part, to newly developed sassays for detecting, diagnosing, monitoring, staging, prognosticating, imaging and treating cancers, particularly breast cancer.

BACKGROUND OF THE INVENTION

- One of every nine American women will develop breast cancer sometime during her life based on a lifespan of 85 years. Annually, over 180,000 women in the United States will be diagnosed with breast cancer and approximately 46,000 will die of the disease.
- Every woman is at risk for breast cancer. A woman's chances of developing breast cancer increase as she grows older; 80 percent of all cancers are found in women over the age of 50. There are also several risk factors that can increase a woman's chances of developing cancer. A woman may 20 be at increased risk if she has a family history of the disease, if she had her first child after the age of 30 or has no children, or if she began menstruating early.

However, more than 70 percent of women who develop breast cancer have no known risk factors. Less than 10 percent of breast cancer cases are thought to be related to the BRCA1 gene discovered in 1994. Researchers are now investigating the role other factors such as nutrition, alcohol, exercise, smoking, and oral contraceptives may play in cancer prevention.

As with many other cancers, the best chance for successful treatment occurs when breast cancer is found early.

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Mammograms, special x-rays of the breast, can detect more than 90 percent of all breast cancers. If breast cancer is found early, the chance of cure is greater than 90 percent. Treatment options include surgery, chemotherapy, and radiation 5 therapy depending on the stage of the cancer.

Procedures used for detecting, diagnosing, monitoring, staging, prognosticating and imaging breast cancer are of critical importance to the outcome of the patient. Patients diagnosed with early breast cancer generally have a much 10 greater five-year survival rate as compared to the survival rate for patients diagnosed with distant metastasized breast cancer. New diagnostic methods which are more sensitive and specific for detecting early breast cancer are clearly needed.

Breast cancer patients are closely monitored following 15 initial therapy and during adjuvant therapy to determine response to therapy and to detect persistent or recurrent disease of metastasis. There is clearly a need for a breast cancer marker which is more sensitive and specific in detecting breast cancer and its recurrence and progression.

Another important step in managing breast cancer is to determine the stage of the patient's disease. determination has potential prognostic value and provides criteria for designing optimal therapy. Generally, pathological staging of breast cancer is preferable over 25 clinical staging because the former gives a more accurate prognosis. However, clinical staging would be preferred were it at least as accurate as pathological staging because it does not depend on an invasive procedure to obtain tissue for pathological evaluation. Staging of breast cancer would be 30 improved by detecting new markers in cells, tissues, or bodily fluids which could differentiate between different stages of invasion.

In the present invention methods are provided for detecting, diagnosing, monitoring, staging, prognosticating, imaging and treating breast cancer via 9 Breast Specific Genes

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(BSGs). The 9 BSGs refer, among other things, to native proteins expressed by the genes comprising the polynucleotide sequences of any of SEQ ID NO: 1-9. In the alternative, what is meant by the 9 BSGs as used herein, means the native mRNAs encoded by the genes comprising any of the polynucleotide sequences of SEQ ID NO: 1-9 or it can refer to the actual genes comprising any of the polynucleotide sequences of SEQ ID NO: 1-9.

Other objects, features, advantages and aspects of the present invention will become apparent to those of skill in the art from the following description. It should be understood, however, that the following description and the specific examples, while indicating preferred embodiments of the invention, are given by way of illustration only. Various changes and modifications within the spirit and scope of the disclosed invention will become readily apparent to those skilled in the art from reading the following description and from reading the other parts of the present disclosure.

SUMMARY OF THE INVENTION

Toward these ends, and others, it is an object of the present invention to provide a method for diagnosing the presence of breast cancer by analyzing for changes in levels of BSG in cells, tissues or bodily fluids compared with levels of BSG in preferably the same cells, tissues, or bodily fluid type of a normal human control, wherein a change in levels of BSG in the patient versus the normal human control is associated with breast cancer.

Further provided is a method of diagnosing metastatic breast cancer in a patient having such cancer which is not 30 known to have metastasized by identifying a human patient suspected of having breast cancer that has metastasized; analyzing a sample of cells, tissues, or bodily fluid from such patient for BSG; comparing the BSG levels in such cells, tissues, or bodily fluid with levels of BSG in preferably the

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same cells, tissues, or bodily fluid type of a normal human control, wherein a change in BSG levels in the patient versus the normal human control is associated with a cancer which has metastasized.

5 Also provided by the invention is a method of staging breast cancer in a human which has such cancer by identifying a human patient having such cancer; analyzing a sample of cells, tissues, or bodily fluid from such patient for BSG; comparing BSG levels in such cells, tissues, or bodily fluid 10 with levels of BSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein a change in BSG levels in the patient versus the normal human control is associated with a cancer which is progressing or regressing or in remission.

Further provided is a method of monitoring breast cancer in a human having such cancer for the onset of metastasis. The method comprises identifying a human patient having such cancer that is not known to have metastasized; periodically analyzing a sample of cells, tissues, or bodily fluid from 20 such patient for BSG; comparing the BSG levels in such cells, tissue, or bodily fluid with levels of BSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein a change in BSG levels in the patient versus the normal human control is associated with a cancer 25 which has metastasized.

Further provided is a method of monitoring the change in stage of breast cancer in a human having such cancer by looking at levels of BSG in a human having such cancer. method comprises identifying a human patient having such 30 cancer; periodically analyzing a sample of cells, tissues, or bodily fluid from such patient for BSG; comparing the BSG levels in such cells, tissue, or bodily fluid with levels of BSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein a change in BSG 35 levels in the patient versus the normal human control is

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associated with a cancer which is progressing or regressing or in remission.

Further provided are antibodies against the BSGs or fragments of such antibodies which can be used to detect or 5 image localization of the BSGs in a patient for the purpose of detecting or diagnosing a disease or condition. antibodies can be polyclonal or monoclonal, or prepared by molecular biology techniques. The term "antibody", as used herein and throughout the instant specification is also meant 10 to include aptamers and single-stranded oligonucleotides such as those derived from an in vitro evolution protocol referred to as SELEX and well known to those skilled in the art. Antibodies can be labeled with a variety of detectable labels including, but not limited to, radioisotopes and paramagnetic 15 metals. These antibodies or fragments thereof can also be used as therapeutic agents in the treatment of diseases characterized by expression of a BSG. In therapeutic applications, the antibody can be used without or with derivatization to a cytotoxic agent such as a radioisotope, 20 enzyme, toxin, drug or a prodrug.

Other objects, features, advantages and aspects of the present invention will become apparent to those of skill in the art from the following description. It should be understood, however, that the following description and the specific examples, while indicating preferred embodiments of the invention, are given by way of illustration only. Various changes and modifications within the spirit and scope of the disclosed invention will become readily apparent to those skilled in the art from reading the following description and from reading the other parts of the present disclosure.

DESCRIPTION OF THE INVENTION

The present invention relates to diagnostic assays and methods, both quantitative and qualitative for detecting, diagnosing, monitoring, staging, prognosticating and imaging

cancers by comparing levels of BSG with those of BSG in a normal human control. What is meant by levels of BSG as used herein, means levels of the native protein expressed by the genes comprising the polynucleotide sequence of any of SEQ ID In the alternative, what is meant by levels of BSG as used herein, means levels of the native mRNA encoded by any of the genes comprising any of the polynucleotide sequences of SEQ ID NO: 1-9 or levels of the gene comprising any of the polynucleotide sequence of SEQ ID NO: 1-9. Such levels are 10 preferably measured in at least one of, cells, tissues and/or bodily fluids, including determination of normal and abnormal levels. Thus, for instance, a diagnostic assay in accordance with the invention for measuring changes in levels of any one of the BSG proteins compared to normal control bodily fluids, 15 cells, or tissue samples may be used to diagnose the presence of cancers, including breast cancer. By "change" it is meant either an increase or decrease in levels of the BSG. For example, for BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) and Mam005 (SEQ ID NO:3), an increase in levels as 20 compared to normal human controls is associated with breast cancer, metastasis and progression of the cancer, while a decrease in levels is association with regression and/or For the BSG Mam002 (SEQ ID NO:1), a decrease in levels as compared to normal human controls is associated with 25 breast cancer, metastasis and progression while an increase is associated with regression and/or remission. Any of the 9 BSGs may be measured alone in the methods of the invention, or all together or any combination of the nine.

All the methods of the present invention may optionally include measuring the levels of other cancer markers as well as BSG. Other cancer markers, in addition to BSG, such as BRCA1 are known to those of skill in the art.

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Diagnostic Assays

The present invention provides methods for diagnosing the presence of breast cancer by analyzing for changes in levels of BSG in cells, tissues or bodily fluids compared with levels of BSG in cells, tissues or bodily fluids of preferably the same type from a normal human control. As demonstrated herein an increase in levels of BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3) in the patient versus the normal human control is associated with the presence of breast cancer, while a decrease in levels of BSGs such as Mam002 (SEQ ID NO:1) in the patient versus the normal human control is associated with the presence of breast cancer.

Without limiting the instant invention, typically, for a quantitative diagnostic assay a positive result indicating the patient being tested has cancer is one in which cells, tissues, or bodily fluid levels of the cancer marker, such as BSG, are at least two times higher or lower, and most preferably are at least five times higher or lower, than in preferably the same cells, tissues, or bodily fluid of a normal human control.

The present invention also provides a method of diagnosing metastatic breast cancer in a patient having breast cancer which has not yet metastasized for the onset of metastasis. In the method of the present invention, a human cancer patient suspected of having breast cancer which may have metastasized (but which was not previously known to have metastasized) is identified. This is accomplished by a variety of means known to those of skill in the art. For example, in the case of breast cancer, patients are typically diagnosed with breast cancer following traditional detection methods.

In the present invention, determining the presence of BSG level in cells, tissues, or bodily fluid, is particularly useful for discriminating between breast cancer which has not

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metastasized and breast cancer which has metastasized. Existing techniques have difficulty discriminating between breast cancer which has metastasized and breast cancer which has not metastasized and proper treatment selection is often dependent upon such knowledge.

In the present invention, the cancer marker levels measured in such cells, tissues, or bodily fluid is BSG, and are compared with levels of BSG in preferably the same cells, tissue, or bodily fluid type of a normal human control. That is, if the cancer marker being observed is just BSG in serum, this level is preferably compared with the level of BSG in serum of a normal human patient. An increase in BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3) in the patient versus the normal human control is associated with breast cancer which has metastasized while a decrease in BSGs such as Mam002 (SEQ ID NO:1) in the patient versus the normal human control is associated with breast cancer which has metastasized.

Without limiting the instant invention, typically, for a quantitative diagnostic assay a positive result indicating the cancer in the patient being tested or monitored has metastasized is one in which cells, tissues, or bodily fluid levels of the cancer marker, such as BSG, are at least two times higher or lower, and most preferably are at least five times higher or lower, than in preferably the same cells, tissues, or bodily fluid of a normal patient.

Normal human control as used herein includes a human patient without cancer and/or non cancerous samples from the patient; in the methods for diagnosing or monitoring for 30 metastasis, normal human control preferably comprises samples from a human patient that is determined by reliable methods to have breast cancer which has not metastasized, such as earlier samples of the same patient.

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Staging

The invention also provides a method of staging breast cancer in a human patient.

The method comprises identifying a human patient having such cancer; analyzing a sample of cells, tissues, or bodily fluid from such patient for BSG. Then, the method compares BSG levels in such cells, tissues, or bodily fluid with levels of BSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein an increase in levels of BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3) or a decrease in levels of BSGs such as Mam002 (SEQ ID NO:1) in the patient versus the normal human control is associated with a cancer which is progressing and a decrease in levels of BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3) or an increase in levels of BSGs such as Mam001 (SEQ ID NO:2) is associated with a cancer which is regressing or in remission.

Monitoring

Further provided is a method of monitoring breast cancer in a human having such cancer for the onset of metastasis. The method comprises identifying a human patient having such cancer that is not known to have metastasized; periodically analyzing a sample of cells, tissues, or bodily fluid from 25 such patient for BSG; comparing the BSG levels in such cells, tissue, or bodily fluid with levels of BSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein an increase in levels of BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3) or a decrease in levels of BSGs such as Mam002 (SEQ ID NO:1) in the patient versus the normal human control is associated with a cancer which has metastasized.

Further provided by this invention is a method of monitoring the change in stage of breast cancer in a human 35 having such cancer. The method comprises identifying a human

patient having such cancer; periodically analyzing a sample of cells, tissues, or bodily fluid from such patient for BSG; comparing the BSG levels in such cells, tissue, or bodily fluid with levels of BSG in preferably the same cells, tissues, or bodily fluid type of a normal human control sample, wherein an increase in levels of BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3) or a decrease in levels of BSGs such as Mam002 (SEQ ID NO:1) in the patient versus the normal human control is associated with a cancer which is progressing in stage and a decrease in the levels of BSGs such as Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3) or an increase in levels of BSGs such as Mam002 (SEQ ID NO:1) is associated with a cancer which is regressing in stage or in remission.

Monitoring such patient for onset of metastasis is periodic and preferably done on a quarterly basis. However, this may be more or less frequent depending on the cancer, the particular patient, and the stage of the cancer.

Assay Techniques

20 Assay techniques that can be used to determine levels of gene expression, such as BSG of the present invention, in a sample derived from a host are well-known to those of skill in the art. Such assay methods include radioimmunoassays, reverse transcriptase PCR (RT-PCR) assays, 25 immunohistochemistry assays, in situ hybridization assays, competitive-binding assays, Western Blot analyses, ELISA assays and proteomic approaches. Among these, ELISAs are frequently preferred to diagnose a gene's expressed protein in biological fluids.

An ELISA assay initially comprises preparing an antibody, if not readily available from a commercial source, specific to BSG, preferably a monoclonal antibody. In addition a reporter antibody generally is prepared which binds specifically to BSG. The reporter antibody is attached to a

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detectable reagent such as radioactive, fluorescent or enzymatic reagent, for example horseradish peroxidase enzyme or alkaline phosphatase.

To carry out the ELISA, antibody specific to BSG is 5 incubated on a solid support, e.g. a polystyrene dish, that binds the antibody. Any free protein binding sites on the dish are then covered by incubating with a non-specific protein such as bovine serum albumin. Next, the sample to be analyzed is incubated in the dish, during which time BSG binds 10 to the specific antibody attached to the polystyrene dish. Unbound sample is washed out with buffer. A reporter antibody specifically directed to BSG and linked to horseradish peroxidase is placed in the dish resulting in binding of the reporter antibody to any monoclonal antibody bound to BSG. 15 Unattached reporter antibody is then washed out. Reagents for peroxidase activity, including a colorimetric substrate are then added to the dish. Immobilized peroxidase, linked to BSG antibodies, produces a colored reaction product. The amount of color developed in a given time period is proportional to 20 the amount of BSG protein present in the sample. Quantitative results typically are obtained by reference to a standard curve.

A competition assay may be employed wherein antibodies specific to BSG attached to a solid support and labeled BSG and a sample derived from the host are passed over the solid support and the amount of label detected attached to the solid support can be correlated to a quantity of BSG in the sample.

Nucleic acid methods may be used to detect BSG mRNA as a marker for breast cancer. Polymerase chain reaction (PCR) and other nucleic acid methods, such as ligase chain reaction (LCR) and nucleic acid sequence based amplification (NASABA), can be used to detect malignant cells for diagnosis and monitoring of various malignancies. For example, reverse-transcriptase PCR (RT-PCR) is a powerful technique which can be used to detect the presence of a specific mRNA population

in a complex mixture of thousands of other mRNA species. In RT-PCR, an mRNA species is first reverse transcribed to complementary DNA (cDNA) with use of the enzyme reverse transcriptase; the cDNA is then amplified as in a standard PCR reaction. RT-PCR can thus reveal by amplification the presence of a single species of mRNA. Accordingly, if the mRNA is highly specific for the cell that produces it, RT-PCR can be used to identify the presence of a specific type of cell.

10 Hybridization to clones or oligonucleotides arrayed on a solid support (i.e., gridding) can be used to both detect the expression of and quantitate the level of expression of that gene. In this approach, a cDNA encoding the BSG gene is fixed to a substrate. The substrate may be of any suitable 15 type including but not limited to glass, nitrocellulose, nylon or plastic. At least a portion of the DNA encoding the BSG gene is attached to the substrate and then incubated with the analyte, which may be RNA or a complementary DNA (cDNA) copy RNA. isolated from the tissue of20 Hybridization between the substrate bound DNA and the analyte can be detected and quantitated by several means including but not limited to radioactive labeling or fluorescence labeling of the analyte or a secondary molecule designed to detect the hybrid. Quantitation of the level of gene expression can be 25 done by comparison of the intensity of the signal from the analyte compared with that determined from known standards. The standards can be obtained by in vitro transcription of the target gene, quantitating the yield, and then using that material to generate a standard curve.

Of the proteomic approaches, 2D electrophoresis is a technique well known to those in the art. Isolation of individual proteins from a sample such as serum is accomplished using sequential separation of proteins by different characteristics usually on polyacrylamide gels.

First, protéins are separated by size using an electric

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current. The current acts uniformly on all proteins, so smaller proteins move farther on the gel than larger proteins. The second dimension applies a current perpendicular to the first and separates proteins not on the basis of size but on the specific electric charge carried by each protein. Since no two proteins with different sequences are identical on the basis of both size and charge, the result of a 2D separation is a square gel in which each protein occupies a unique spot. Analysis of the spots with chemical or antibody probes, or subsequent protein microsequencing can reveal the relative abundance of a given protein and the identity of the proteins in the sample.

The above tests can be carried out on samples derived from a variety of patients' cells, bodily fluids and/or tissue extracts (homogenates or solubilized tissue) such as from tissue biopsy and autopsy material. Bodily fluids useful in the present invention include blood, urine, saliva, or any other bodily secretion or derivative thereof. Blood can include whole blood, plasma, serum, or any derivative of blood.

In Vivo Antibody Use

Antibodies against BSGs can also be used *in vivo* in patients with disease of the breast. Specifically, antibodies against a BSG can be injected into a patient suspected of 25 having a disease of the breast for diagnostic and/or therapeutic purposes. The use of antibodies for *in vivo* diagnosis is well known in the art. For example, antibody-chelators labeled with Indium-111 have been described for use in the radioimmunoscintographic imaging of carcinoembryonic antigen expressing tumors (Sumerdon et al. Nucl. Med. Biol. 1990 17:247-254). In particular, these antibody-chelators have been used in detecting tumors in patients suspected of having recurrent colorectal cancer (Griffin et al. J. Clin. Onc. 1991 9:631-640). Antibodies with paramagnetic ions as

labels for use in magnetic resonance imaging have also been described (Lauffer, R.B. Magnetic Resonance in Medicine 1991 22:339-342). Antibodies directed against BSGs can be used in a similar manner. Labeled antibodies against a BSG can be 5 injected into patients suspected of having a disease of the breast such as breast cancer for the purpose of diagnosing or staging of the disease status of the patient. The label used will be selected in accordance with the imaging modality to be used. For example, radioactive labels such as Indium-111, 10 Technetium-99m or Iodine-131 can be used for planar scans or single photon emission computed tomography (SPECT). emitting labels such as Fluorine-19 can be used in positron emission tomography. Paramagnetic ions such as Gadlinium (III) or Manganese (II) can used in magnetic resonance imaging Localization of the label within the breast or external to the breast permits determination of the spread of the disease. The amount of label within the breast also allows determination of the presence or absence of cancer in the breast.

20 For patients diagnosed with breast cancer, injection of an antibody against a BSG can also have a therapeutic benefit. The antibody may exert its therapeutic effect alone. Alternatively, the antibody is conjugated to a cytotoxic agent such as a drug, toxin or radionuclide to enhance its 25 therapeutic effect. Drug monoclonal antibodies have been described in the art for example by Garnett and Baldwin, Cancer Research 1986 46:2407-2412. The use of toxins conjugated to monoclonal antibodies for the therapy of various cancers has also been described by Pastan et al. Cell 1986 30 47:641-648). Yttrium-90 labeled monoclonal antibodies have been described for maximization of dose delivered to the tumor while limiting toxicity to normal tissues (Goodwin and Meares Cancer Supplement 1997 80:2675-2680). Other cytotoxic radionuclides including, but not limited to Copper-67, Iodine-

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131 and Rhenium-186 can also be used for labeling of antibodies against BSGs.

Antibodies which can be used in these in vivo methods include both polyclonal and monoclonal antibodies 5 antibodies prepared via molecular biology techniques. Antibody fraqments and aptamers and single-stranded oligonucleotides such as those derived from an in vitro evolution protocol referred to as SELEX and well known to those skilled in the art can also be used.

10 EXAMPLES

The present invention is further described by the following examples. The examples are provided solely to illustrate the invention by reference to specific embodiments. These exemplifications, while illustrating certain specific aspects of the invention, do not portray the limitations or circumscribe the scope of the disclosed invention.

Example 1

Identification of BSGs were carried out by a systematic analysis of data in the LIFESEQ database available from Incyte 20 Pharmaceuticals, Palo Alto, CA, using the data mining Cancer Leads Automatic Search Package (CLASP) developed by diaDexus LLC, Santa Clara, CA.

The CLASP performs the following steps:

Selection of highly expressed organ specific genes based 25 on the abundance level of the corresponding EST in the targeted organ versus all the other organs.

Analysis of the expression level of each highly expressed organ specific genes in normal, tumor tissue, disease tissue and tissue libraries associated with tumor or 30 disease.

Selection of the candidates demonstrating component ESTs were exclusively or more frequently found in tumor libraries.

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CLASP allows the identification of highly expressed organ and cancer specific genes useful in the diagnosis of breast cancer.

Table 1: BSGs Sequences

5	SEQ ID NO:	LS Clone ID	LSA Gene ID
	. 1	2740238 (Mam002)	242151
	2	1730886 (Mam001)	238469
	3	y155b03 (Mam005)	348845
	4	2613064 (Mam004)	27052
10	5	894184	221086
	6	2299454	27681
	7	2258254	248176
	8	789767	156580
•	9	1213903	219737

The following example was carried out using standard techniques, which are well known and routine to those of skill in the art, except where otherwise described in detail. Routine molecular biology techniques of the following example can be carried out as described in standard laboratory manuals, such as Sambrook et al., MOLECULAR CLONING: A LABORATORY MANUAL, 2nd Ed.; Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (1989).

Example 2: Relative Quantitation of Gene Expression

PCR with fluorescent Taqman probes is a quantitative detection system utilizing the 5'-3' nuclease activity of Taq DNA polymerase. The method uses an internal fluorescent oligonucleotide probe (Taqman) labeled with a 5' reporter dye and a downstream, 3' quencher dye. During PCR, the 5'-3' nuclease activity of Taq DNA polymerase releases the reporter, whose fluorescence can then be detected by the laser detector of the Model 7700 Sequence Detection System (PE Applied Biosystems, Foster City, CA, USA).

Amplification of an endogenous control was used to standardize the amount of sample RNA added to the reaction and normalize for Reverse Transcriptase (RT) efficiency. cyclophilin, glyceraldehyde-3-phosphate dehydrogenase (GAPDH) 5 or 18S ribosomal RNA (rRNA) was used as this endogenous control. To calculate relative Quantitation between all the samples studied, the target RNA levels for one sample were used as the basis for comparative results (calibrator). Quantitation relative to the "calibrator" can be obtained 10 using the standard curve method or the comparative method (User Bulletin #2: ABI PRISM 7700 Sequence Detection System). To evaluate the tissue distribution, and the level of breast specific markers (BSM) Mam001 (SEQ ID NO:2), Mam002 (SEQ ID NO:1), Mam004 (SEQ ID NO:4) and Mam005 (SEQ ID NO:3) in normal 15 and cancer tissue, total RNA was extracted from cancer and matched normal adjacent tissues (NAT) and from unmatched cancer and normal tissues. Subsequently, first strand cDNA was prepared with reverse transcriptase and the polymerase chain reaction carried out using primers and Tagman probes 20 specific to each of Mam001 (SEQ ID NO:2), Mam002 (SEQ ID Mam004 (SEQ ID NO:4) and Mam005 (SEQ ID respectively. The results are obtained using the ABI PRISM 7700 Sequence Detector. The numbers are relative levels of expression of Mam001 (SEQ ID NO:2), Mam002 (SEQ ID NO:1), 25 Mam004 (SEQ ID NO:4) and Mam005 (SEQ ID NO:3) compared to their respective calibrators.

Measurement of SEQ ID NO:2; Clone ID:1730886; Gene ID: 238469 (Mam001)

The numbers depicted in Table 2 are relative levels of 30 expression in 12 normal tissues of Mam001 (SEQ ID NO:2) compared to testis (calibrator). These RNA samples were obtained commercially and were generated by pooling samples from a particular tissue from different individuals.

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Table 2: Relative levels of Mam001 (SEQ ID NO:2) Expression in Pooled Samples

Tissue	NORMAL
Brain	0
Heart	0
Kidney	0
Liver	0
Lung	0
Mammary	6
Prostate	0
Muscle	0
Small Intestine	0
Testis	1
Thymus	0
Uterus	0

The relative levels of expression in Table 2 show that Mam001 (SEQ ID NO:2) mRNA expression is detected in the pool of normal mammary and in testis but not in the other 10 normal tissue pools analyzed. These results demonstrate that Mam001 (SEQ ID NO:2) mRNA expression is highly specific for mammary tissue and is also found in testis. Expression in a male specific tissue is not relevant in detecting cancer in female specific tissues

The tissues shown in Table 2 are pooled samples from different individuals. The tissues shown in Table 3 were obtained from individuals and are not pooled. Hence the values for mRNA expression levels shown in Table 2 cannot be directly compared to the values shown in Table 3.

The numbers depicted in Table 3 are relative levels of 30 expression of Mam001 (SEQ ID NO:2) compared to testis (calibrator), in 24 pairs of matching samples. Each matching pair contains the cancer sample for a particular tissue and the normal adjacent tissue (NAT) sample for that same tissue from the same individual.

Table 3: Relative levels of Mam001 (SEQ ID NO:2) Expression in Individual Samples

	Sample ID	Tissue	Cancer	Matching Normal
	Mam 47XP	Mammary Gland	0	0
5	Mam A06X	Mammary Gland	23.	1
	Mam B011X	Mammary Gland	0	5
	Mam 603X/C034	Mammary Gland	0	2.10
	Mam 162X	Mammary Gland	1.96	0.15
	Mam 42DN	Mammary Gland	0.38	1.27
10	Mam S079	Mammary Gland	0.34	0.36
	Mam S123	Mammary Gland	0.03	0.87
	Mam S516	Mammary Gland	0.43	0.53
	Mam S699	Mammary Gland	0.40	0.66
	Mam S997	Mammary Gland	0.41	0.51
15	Sto AC44	Stomach	0	0
	TST 39X	Testis	0	0
	Cln SG45	Colon	0	0
	Cln TX01	Colon	0	0
	Cvx NK23	Cervix	0	0
20	Cvx NK24	Cervix	0	0
	Endo 3AX	Endometrium	0	0
	Endo 4XA	Endometrium	0	0
	Endo 5XA	Endometrium	0	0
:	Kid 11XD	Kidney	0	0
25	Kid 5XD	Kidney	0	0
	Lng C20X	Lung	0	0
	Lng SQ56	Lung	0	0

Among 48 samples in Table 3 representing 8 different tissues expression is seen only in mammary tissues. These 30 results confirm the tissue specificity results obtained with

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normal samples shown in Table 2. Table 2 and Table 3 represent a combined total of 60 samples in 16 human tissue types. Thirty-six samples representing 14 different tissue types excluding breast and testis had no detected Mam001 (SEQ ID NO:2) mRNA (Table 2 and 3). Other than breast tissue, Mam001 (SEQ ID NO:2) is detected only in one other tissue type (Testis) and then only in the pooled tissue sample (Table 2) but not in the matched testis cancer samples (Table 3).

Comparisons of the level of mRNA expression in breast 10 cancer samples and the normal adjacent tissue from the same individuals are shown in Table 3. Mam001 (SEQ ID NO:2) is expressed at higher levels in 2 of 11 breast cancer tissues (Mam A06X and Mam 162X) compared with the corresponding normal adjacent tissue. The level of Mam001 (SEQ ID NO:2) expression 15 is lower in breast cancer compared to normal adjacent tissue in four matched samples (Mam B011X, Mam 603X/CO34, Mam 42DN No expression was detected in one set of and Mam S123). matched samples (Mam 47XP). Equivalent levels or very similar levels of expression were detected in four other matched 20 samples (Mam S079, Mam S516, Mam S699 and Mam S997). increasing cancer mass might in these cases result in an overall increase in the total amount of expression.

The high level of tissue specificity and increased or equivalent expression in 6 of 11 individuals is demonstrative 25 of Mam001 (SEQ ID NO:2) being a diagnostic marker for detection of mammary cancer cells using mRNA.

Measurement of SEQ ID NO:1; Clone ID: 2740238; Gene ID 242151 (Mam002)

The numbers depicted in Table 5 are relative levels of 30 expression in 12 normal tissues of Mam002 (SEQ ID NO:1) compared to Thymus (calibrator). These RNA samples were obtained commercially and were generated by pooling samples from a particular tissue from different individuals.

Table 4: Relative levels of Mam002 (SEQ ID NO:1) Expression in Pooled Samples

	Tissue	NORMAL
	Brain	0.03
5	Heart	0.01
	Kidney	0
	Liver	0
	Lung	0.06
	Mammary	289.01
10	Muscle	0
	Prostate	0.31
	Small Int.	0
	Testis	0.08
	Thymus	1.00
15	Uterus	0

The relative levels of expression in Table 4 show that Mam002 (SEQ ID NO:1) mRNA expression is detected at a high level in the pool of normal mammary but at very low levels in the other 11 normal tissue pools analyzed. These results demonstrate that Mam002 (SEQ ID NO:1) mRNA expression is highly specific for mammary tissue.

The tissues shown in Table 4 are pooled samples from different individuals. The tissues shown in Table 5 were obtained from individuals and are not pooled. Hence the values for mRNA expression levels shown in Table 4 cannot be directly compared to the values shown in Table 5.

The numbers depicted in Table 5 are relative levels of expression of Mam002 (SEQ ID NO:1) compared to thymus (calibrator) in 27 pairs of matching samples. Each matching 30 pair contains the cancer sample for a particular tissue and the normal adjacent tissue (NAT) sample for that same tissue from the same individual. In addition 2 unmatched mammary samples from normal tissues and one unmatched ovarian cancer and one normal (non-cancerous) ovary were also tested.

Table 5: Relative levels of Mam002 (SEQ ID NO:1) Expression in Individual Samples

	Sample ID	Tissue	Cancer	Matching	Normal
	Mam 12X	Mammary Gland	7.2	69	
5	Mam 42DN	Mammary Gland	1051	. 2075	
	Mam 59X	Mammary Gland	7.0	15.5	
	Mam A06X	Mammary Gland	1655	1781	
	Mam B011X	Mammary Gland	32.1	2311	
	Mam S127	Mammary Gland	1.73	0	
10	Mam S516	Mammary Gland	9.72	69.95	
	Mam S699	Mammary Gland	83.46	75.65	
	Mam S854	Mammary Gland	133.23	836.56	
	Mam S967	Mammary Gland	59.77	188.28	·
	Mam S997	Mammary Gland	94.14	73.64	
15	Mam 162X	Mammary Gland	674.0	31.1	
	Mam C012	Mammary Gland	N/A	N/A	11379.3
	Mam C034	Mammary Gland	N/A	N/A	3502.6
	Mam S079	Mammary Gland	11772.5	903.5	
	Mam S123	Mammary Gland	3.4	170.5	
20	Ovr 103X	Ovary	0	0	
	Ovr 1118	Ovary	0.13	N/A	<u> </u>

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	Ovr 35GA	Ovary	N/A	N/A	0.13
	Utr 23XU	Uterus	5.6	0	
	Utr 135X0	Uterus	0	0	
	Cvx NK24	Cervix	0.9	1.4	
5	End 4XA	Endometriu m	32.2	0	
	Cln AS43	Colon	2.3	0	
	Cln AS45	Colon	. 0	0	
	Cln RC01	Colon	0.2	0	
	Lng AC90	Lung	0	2.0	
10	Lng LC109	Lung	0	0.6	
	Lng SC32	Lung	0.8	0	
	Sto AC93	Stomach	0	0	
-	Tst 39X	Testis	1.97	0	

Among 58 samples in Table 5 representing 9 different tissues, the highest expression is seen in mammary tissues. Amongst the non-breast tissues which show expression, only one sample (End 4XA) has expression comparable to that seen in the majority of the breast samples tested. This sample is endometrial tissue, which is a female specific tissue. These results confirm the tissue specificity results obtained with normal samples shown in Table 4. Table 4 and Table 5 represent a combined total of 70 samples in 17 human tissue types. Twenty-two samples representing 11 different tissue types excluding breast had no detected Mam002 (SEQ ID NO:1) mRNA (Table 4 and Table 5).

Comparisons of the level of mRNA expression in breast cancer samples and the normal adjacent tissue from the same individuals are shown in Table 5. Mam002 (SEQ ID NO:1) is expressed at higher levels in 3 of 13 matched breast cancer tissues (Samples Mam S127, Mam 162X and Mam S079) compared with the corresponding normal adjacent tissue. The level of Mam002 (SEQ ID NO:1) expression is lower in breast cancer

compared to normal adjacent tissue in eight individuals (Mam 12X, Mam 42DN, Mam 59X, Mam B011X, Mam S516, Mam S854, Mam S967, and Mam S123). Equivalent levels or very similar levels of expression were detected in three other matched samples 5 (Samples Mam A06X, Mam S699 and Mam S997).

The high level of tissue specificity is demonstrative of Mam002 (SEO ID NO:1) being a diagnostic marker for detection of mammary cancer cells using mRNA. Breast tissue is the only significant source of this gene's expression so 10 far detected. Eight of 13 matched samples have lower levels of expression in cancer than normal adjacent tissue. decreased expression of this gene appears to be diagnostic of cancer presence.

Measurement of SEQ ID NO:4; Clone ID: 2613064; Gene ID: 27052 15 (Mam004)

The numbers depicted in Table 6 are relative levels of expression in 12 normal tissues of Mam004 (SEQ ID NO:4) compared to mammary (calibrator). These RNA samples were obtained commercially and were generated by pooling samples 20 from a particular tissue from different individuals.

Relative levels of Mam004 (SEQ ID NO:4) Expression in Pooled Samples

	Tissue	NORMAL
	Brain	0.059
25	Heart	0.131
	Kidney	0.018
	Liver	0
	Lung	0.478
	Mammary	1.000
30	Prostate	0.459
	Muscle	0.003
	Small Intestine	0.048
	Testis	0.130
	Thymus	0.030
35	Uterus	0.071

The relative levels of expression in Table 6 show that Mam004 (SEQ ID NO:4) mRNA expression is detected in the pool of

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normal mammary and also in other tissues including lung, prostate, testis and heart. These results demonstrate that although more highly expressed in normal breast tissue Mam004(SEQ ID NO:4) mRNA expression is not specific for mammary gland.

The tissues shown in Table 6 are pooled samples from different individuals. The tissues shown in Table 7 were obtained from individuals and are not pooled. Hence the values for mRNA expression levels shown in Table 6 cannot be directly compared to the values shown in Table 7.

The numbers depicted in Table 7 are relative levels of expression of Mam004 (SEQ ID NO:4) compared to mammary (calibrator), in 23 pairs of matching samples. Each matching pair contains the cancer sample for a particular tissue and the normal adjacent tissue (NAT) sample for that same tissue from the same individual.

Table 7: Relative levels of Mam004 (SEQ ID NO:4) Expression in Individual Samples

	Sample ID	Tissue	Cancer	Matching
20	Mam 12B	Mammary Gland	0	0
	Mam 12X	Mammary Gland	13.454	0
	Mam 603X	Mammary Gland	30.484	0
	Mam 59X	Mammary Gland	1.306	0
	Mam 162X	Mammary Gland	0.71	0.04
25	Mam 42DN	Mammary Gland	0.25	2.17
	Mam S079	Mammary Gland	42.18	0.47
	Mam S123	Mammary Gland	0.01	0
	Mam S516	Mammary Gland	1.17	0.41
	Mam S699	Mammary Gland	0.11	0.55
30	Mam S997	Mammary Gland	10.43	1.29
	Sto AC44	Stomach	0.61	0

Cln SG45		Colon	0.04	0
	Cln TX01	Colon	0	0
	Cvx NK23	Cervix	0	0
i	Cvx NK24	Cervix	0	0
5	Endo 3AX	Endometrium	0	0
	Endo 4XA	Endometrium	0	0
	Endo 5XA	Endometrium	0	2.73
	Kid 11XD	Kidney	0	0
	Kid 5XD	Kidney	0	2.63
10	Lng C20X	Lung	0	0
	Lng SQ56	Lung	10.37	0

Among 46 samples in Table 7 representing 7 different tissues expression is highest in breast tissues particularly cancers. Expression comparable to that seen in breast samples is also seen in 1 of 4 lung samples (Sample 23), 1 of 4 kidney samples (Sample 21) and 1 of 6 endometrial samples (Sample 19). Table 6 and Table 7 represent a combined total of 58 samples in 16 human tissue types. Twenty samples representing 7 different tissue types excluding breast had no detected 20 Mam004 (SEQ ID NO:4) mRNA (Table 6 and Table 7).

Comparisons of the level of mRNA expression in breast cancer samples and the normal adjacent tissue from the same individuals are shown in Table 7. Mam004 (SEQ ID NO:4) is expressed at higher levels in 8 of 11 breast cancer tissues (Mam 12X, Mam 603X, Mam 59X, Mam 162X, Mam 5079, Mam 5123, Mam 5516 and Mam S997) compared with the corresponding normal adjacent tissue. The level of Mam004 (SEQ ID NO:4) expression is lower in breast cancer compared to normal adjacent tissue in two matched samples (Mam 42DN and Mam S699). No expression was detected in one matched sample (Mam 12B).

Elevated expression in the majority of matched cancer samples compared to normal adjacent tissue is indicative of

Mam004 (SEQ ID NO:4) being a diagnostic marker for detection of mammary cancer cells using mRNA.

Measurement of SEQ ID NO:3; Clone ID:y155b03; Gene ID: 348845 (Mam005)

The numbers depicted in Table 8 are relative levels of expression in 12 normal tissues of Mam005 (SEQ ID NO:3) compared to testis (calibrator). These RNA samples were obtained commercially and were generated by pooling samples from a particular tissue from different individuals.

10 Table 8: Relative levels of Mam005 (SEQ ID NO:3) Expression in Pooled Samples

Tissue	NORMAL
Brain	0
Heart	0.0002
Kidney	0.0001
Liver	0
Lung	0
Mammary	5.4076
Muscle	0
Prostate	0
Small Intestine	0
Testis	1
Thymus	0
Uterus	0

The relative levels of expression in Table 8 show that Mam005 (SEQ ID NO:3) mRNA expression is detected in the pool of normal mammary and in testis but is not present at significant levels in the other 10 normal tissue pools analyzed. These results demonstrate that Mam005 (SEQ ID NO:3) mRNA expression is highly specific for mammary tissue and is also found in testis. Expression in a male specific tissue is not relevant in detecting cancer in female specific tissues.

The tissues shown in Table 8 are pooled samples from different individuals. The tissues shown in Table 9 were 35 obtained from individuals and are not pooled. Hence the values for mRNA expression levels shown in Table 8 cannot be directly compared to the values shown in Table 9.

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The numbers depicted in Table 9 are relative levels of expression of Mam005 (SEQ ID NO:3) compared to testis (calibrator), in 46 pairs of matching samples. Each matching pair contains the cancer sample for a particular tissue and 5 the normal adjacent tissue sample for that same tissue from the same individual. In addition 2 unmatched mammary samples from normal tissues and one unmatched ovarian cancer and one normal (non-cancerous) ovary were also tested.

Table 9: Relative levels of Mam005 (SEQ ID NO:3) Expression 10 in Individual Samples

	Sample ID	Tissue	Cancer	Matching	Normal
	Mam 12X	Mammary Gland	0.33	0.71	
	Mam 42DN	Mammary Gland	0.22	0.63	
	Mam 59X	Mammary Gland	0.03	0.23	
15	Mam A06X	Mammary Gland	70.77	0.56	
	Mam B011X	Mammary Gland	0.03	1.52	
	Mam 162X	Mammary Gland	0.43	0.09	
	Mam C012	Mammary Gland	N/A	N/A	1.6
	Mam C034	Mammary Gland	N/A	N/A	2.9
20	Mam S079	Mammary Gland	0.22	0.13	
	Mam S123	Mammary Gland	0.01	0.23	
	Mam S127	Mammary Gland	0	0.28	
	Mam S516	Mammary Gland	0.15	0.05	

	Mam S699	Mammary Gland	0.21	0.42	
	Mam S854	Mammary Gland	1.12	0.54	
	Mam S967	Mammary Gland	30.61	0.54	
	Mam S997	Mammary Gland	0.40	0.22	
5	Mam 14DN	Mammary Gland	0.07	0	
	Mam 699F	Mammary Gland	0.01	0.09	
	Mam S621	Mammary Gland	1.82	0	
	Mam S918	Mammary Gland	6.89	1.06	·
	Cln CM67	Colon	0	0	
10	Cln DC19	Colon	0	0	
	Cln AS43	Colon	0	0	
	Cln AS45	Colon	0	0	
	Cln RC01	Colon	0.0012	0.0003	
	Lng AC90	Lung	0	0	
15	Lng LC109	Lung	0	0	
	Lng SQ32	Lung	0	0	
	Lng SQ43	Lung	0	0	
	Ovr 103X	Ovary	0	0	
	Ovr 1118	Ovary	0	N/A	
20	Ovr A084	Ovary	0	0	
:	Ovr G021	Ovary	0	0	
	Ovr 35GA	Ovary	N/A	N/A	0
	Cvx NK23	Cervix	0	0	
	Cvx NK24	Cervix	0	0	
25	Endo 3AX	Endometriu m	0	0	

	Endo 4XA	Endometriu m	0	0	
	Sto 758S	Stomach	0	0	
	Sto AC44	Stomach	0	0	
	Sto AC93	Stomach	0	0	
5	Tst 39X	Testis	0.01	0.01	
	Utr 85XU	Uterus	0	0	
	Utr 135XO	Uterus	0	0	
	Utr 23XU	Uterus	0	0	
	Kid 124D	Kidney	0	0	
10	Lvr 15XA	Liver	0	0	
	Pan CO44	Pancreas	0	0	
	Skn 448S	Skin	0	0	
	SmInt 21XA	Small Intestines	0	0	

Among 96 samples in Table 9 representing 14 different tissues significant expression is seen only in breast tissues. These results confirm the tissue specificity results obtained with normal samples shown in Table 8. Table 8 and Table 9 represent a combined total of 108 samples in 18 human tissue types. Sixty-seven samples representing 16 different tissue types excluding breast and testis had either no or very low levels of detected Mam005 (SEQ ID NO:3) mRNA (Table 8 and Table 9).

Comparisons of the level of mRNA expression in breast cancer samples and the normal adjacent tissue from the same individuals are shown in Table 9. Mam005 (SEQ ID NO:3) is expressed at higher levels in 10 of 18 cancer and normal adjacent tissue samples (Mam A06X, Mam 162X, Mam S079, Mam S516, Mam S854, Mam S967, Mam S997, Mam 14DN, Mam S621, and Mam S918) compared with the corresponding normal adjacent tissue. The level of Mam005 (SEQ ID NO:3) expression is lower in breast cancer compared to normal adjacent tissue in eight

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cancer and normal adjacent tissue samples (Mam 12X, Mam 42DN, Mam 59X, Mam B011X, Mam S123, Mam S127, Mam S699 and Mam 699F). No expression was detected in two matching samples.

The high level of tissue specificity and overexpression in 10 of 18 matched cancer and normal adjacent tissue samples is indicative of Mam005 (SEQ ID NO:3) being a diagnostic marker for detection of mammary cancer cells using mRNA.

What is claimed is:

- 1. A method for diagnosing the presence of breast cancer in a patient comprising:
- (a) measuring levels of BSG in cells, tissues or bodily 5 fluids in said patient; and
- (b) comparing measured levels of BSG with levels of BSG in cells, tissues or bodily fluids from a normal human control, wherein a change in measured levels of BSG in the patient versus normal human control is associated with the 10 presence of breast cancer.
 - 2. A method of diagnosing metastatic breast cancer in a patient having breast cancer comprising:
 - (a) identifying a patient having breast cancer that is not known to have metastasized;
- 15 (b) measuring levels of BSG in a sample of cells, tissues, or bodily fluid from said patient; and
- (c) comparing the measured BSG levels with levels of BSG in cells, tissue, or bodily fluid type of a normal human control, wherein a change in measured BSG levels in the patient versus the normal human control is associated with a cancer which has metastasized.
 - 3. A method of staging breast cancer in a patient comprising:
 - (a) identifying a patient having breast cancer;
 - (b) measuring levels of BSG in a sample of cells, tissues, or bodily fluid from said patient for BSG; and
- (c) comparing measured BSG levels with levels of BSG in cells, tissues, or bodily fluid type of a normal human control sample, wherein a change in measured BSG levels in said patient versus the normal human control is associated with a cancer which is progressing or regressing or in remission.

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- 4. A method of monitoring breast cancer in a patient having breast cancer for the onset of metastasis comprising:
- (a) identifying a patient having breast cancer that is not known to have metastasized;
- 5 (b) periodically measuring BSG levels in a sample of cells, tissues, or bodily fluid from said patient; and
- (c) comparing the measured BSG levels with levels of BSG in cells, tissues, or bodily fluid type of a normal human control, wherein a change in BSG levels in the patient versus the normal human control is associated with a cancer which has metastasized.
 - 5. A method of monitoring the change in stage of breast cancer in a patient having breast cancer comprising:
 - (a) identifying a patient having breast cancer;
- (b) periodically measuring BSG levels in a sample of cells, tissues, or bodily fluid from said patient; and
- (c) comparing the measured BSG levels with levels of BSG in cells, tissues, or bodily fluid type of a normal human control, wherein a change in measured BSG levels in the patient versus the normal human control is associated with a cancer which is progressing in stage, which is regressing in stage, or in remission.
- 6. The method of claim 1, 2, 3, 4 or 5 wherein the change associated with the presence, metastasis or progression of breast cancer in said patient is an increase in measured BSG levels in the patient and the BSG comprises Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3).
- 7. The method of claim 1, 2, 3, 4 or 5 wherein the change associated with the presence, metastasis or progression 30 of breast cancer in said patient is a decrease in measured BSG levels in the patient and the BSG comprises Mam002 (SEQ ID NO:1).

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- 8. The method of claim 3 or 5 wherein the change associated with the regression or remission of breast cancer in said patient is a decrease in measured BSG levels in the patient and the BSG comprises Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3).
 - 9. The method of claim 3 or 5 wherein the change associated with the regression or remission of breast cancer in said patient is an increase in measured BSG levels in the patient and the BSG comprises Mam002 (SEQ ID NO:1).
- 10 10. An antibody against a BSG wherein said BSG comprises Mam001 (SEQ ID NO:2), Mam004 (SEQ ID NO:4) or Mam005 (SEQ ID NO:3).
- 11. A method of imaging breast cancer in a patient comprising administering to the patient an antibody of claim 15 10.
 - 12. The method of claim 11 wherein said antibody is labeled with paramagnetic ions or a radioisotope.
- 13. A method of treating breast cancer in a patient comprising administering to the patient an antibody of claim 20 10.
 - 14. The method of claim 13 wherein the antibody is conjugated to a cytotoxic agent.

SEQUENCE LISTING

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/16811

A. CLASSIFICATION OF SUBJECT MATTER					
\ - >	:Please See Extra Sheet. :435/6, 7.1, 91.2; 536/23.5, 24.31; 424/174.1; 530	/388.1. 388.8			
\$	o International Patent Classification (IPC) or to both				
B. FIEL	DS SEARCHED				
Minimum d	ocumentation searched (classification system followe	d by classification symbols)	- ""		
U.S. :	435/6, 7.1, 91.2; 536/23.5, 24.31; 424/174.1; 530/	388.1, 388.8			
Documentat	ion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched		
	lata base consulted during the international search (nee Extra Sheet.	ame of data base and, where practicable	e, search terms used)		
c. Doc	UMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.		
х	US 5,668,267 A (WATSON et al) 16 and 7.	1-5			
Y	WO 98/18945 A (ABBOTT LABORA) 4, 7, 46 and 87.	1-5, 7, 9			
x	US 5,759,776 A (SMITH et al) 02 June 1998, column 16.				
Purther documents are listed in the continuation of Box C. See patent family annex.					
Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand					
	cument defining the general state of the art which is not considered be of particular relevance	the principle or theory underlying the			
"B" car	tier document published on or after the international filing date	"X" document of particular relevance; the			
cit	cument which may throw doubts on priority claim(s) or which is ed to establish the publication date of another citation or other social reason (as specified)	"Y" document of particular relevance; the	e claimed invention cannot be		
	cument referring to an oral disclosure, use, exhibition or other	considered to involve an inventive combined with one or more other sucl being obvious to a person skilled in t	h documents, such combination		
P document published prior to the international filing date but later than the priority date claimed *& document member of the same patent family					
Date of the actual completion of the international search 21 SEPTEMBER 1999 Date of mailing of the international search report 20 OOT 1999					
21 SEPTE	EMBER 1999	20 001 1000			
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/16811

A. CLA	SSIFICATION	OF	SUBJECT	MATTER:
IPC (6):	:			

C12Q 1/68; C12P 19/34; C07H 21/04; A61K 16/00; G01N 33/53

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS; WEST Derwent files; DIALOG: Medline, Biosis, Embase, Scisearach, CA; GenBank/EMBL, n-geneseq search terms: breast, mammary, tumor, carcinoma, cancer, mRNA, protein, antibody, SEQ ID NO: 1-5

Form PCT/ISA/210 (extra sheet)(July 1992)★

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